

112TH CONGRESS
2D SESSION

H. R. 5866

To enhance Food and Drug Administration oversight of medical device recalls, to provide for the conditional clearance of certain medical devices, and for other purposes.

IN THE HOUSE OF REPRESENTATIVES

MAY 30, 2012

Mr. BRALEY of Iowa introduced the following bill; which was referred to the Committee on Energy and Commerce

A BILL

To enhance Food and Drug Administration oversight of medical device recalls, to provide for the conditional clearance of certain medical devices, and for other purposes.

Be it enacted by the Senate and House of Representatives of the United States of America in Congress assembled,

3 SECTION 1. SHORT TITLE.

4 This Act may be cited as the “Medical Device Patient
5 Safety Act”.

6 SEC. 2. OVERSIGHT OF DEVICE RECALLS BY THE FOOD

AND DRUG ADMINISTRATION

8 (a) DEFINITIONS.—In this Act:

1 (1) COMMISSIONER.—The term “Commissioner” means the Commissioner of Food and
2 Drugs.

3 (2) DEVICE.—The term “device” has the meaning given that term in section 201(h) of the Federal
4 Food, Drug, and Cosmetic Act (21 U.S.C. 321(h)).

5 (3) SECRETARY.—The term “Secretary” means
6 the Secretary of Health and Human Services.

7 (b) ACTIONS BY SECRETARY.—To enhance the oversight by the Food and Drug Administration of device recalls, the Secretary of Health and Human Services, acting through the Commissioner of Food and Drugs, shall carry out the activities described in this section.

8 (c) ASSESSMENT OF DEVICE RECALL INFORMATION.—

9 (1) IN GENERAL.—

10 (A) ASSESSMENT PROGRAM.—The Secretary shall establish a program to routinely and systematically assess—

11 (i) information submitted to the Secretary pursuant to a device recall order under section 518(e) of the Federal Food, Drug, and Cosmetic Act (21 U.S.C. 360h(e)); and

(ii) information required to be reported to the Secretary regarding a correction or removal of a device under section 519(g) of such Act (21 U.S.C. 360i(g)).

(B) USE.—The Secretary shall use the assessment of information described under subparagraph (A) to proactively identify strategies for mitigating health risks presented by defective or unsafe devices.

(2) DESIGN.—The program under paragraph

(1) shall be designed, at a minimum, to identify—

(A) trends in the numbers and types of de-

vice recalls;

(B) the types of devices in each device

class that are most frequently recalled;

(C) the causes of device recalls;

(D) the length of time needed for a person

subject to a device recall to complete the recall;

(E) the length of time needed

retary to terminate a device recall;

(F) whether the Secretary has

device recall audit check;

(G) which persons h

1 (H) any other information as the Secretary
2 determines appropriate.

3 (d) AUDIT CHECK PROCEDURES.—The Secretary
4 shall clarify procedures for conducting device recall audit
5 checks to improve the ability of investigators to perform
6 these checks in a consistent manner.

7 (e) ASSESSMENT CRITERIA.—The Secretary shall de-
8 velop explicit criteria for assessing whether a person sub-
9 ject to a recall order under section 518(e) of the Federal
10 Food, Drug, and Cosmetic Act (21 U.S.C. 360h(e)) or to
11 a requirement under section 519(g) of such Act (21
12 U.S.C. 360i(g)) has performed an effective correction or
13 removal action under such section 519(g).

14 (f) TERMINATION OF RECALLS.—

15 (1) IN GENERAL.—The Secretary shall docu-
16 ment the basis for the termination by the Food and
17 Drug Administration of—

18 (A) an individual device recall ordered
19 under section 518(e) of the Federal Food,
20 Drug, and Cosmetic Act (21 U.S.C. 360h(e));
21 and

22 (B) the requirement on a manufacturer or
23 importer of a device to report any correction or
24 removal action for which a report is required to

1 be submitted to the Secretary under section
2 519(g) of such Act (21 U.S.C. 360i(g)).

3 (2) PUBLICATION.—

4 (A) IN GENERAL.—The Secretary shall,
5 with respect to each termination described in
6 paragraph (1), publish the documentation re-
7 quired under such paragraph not later than 180
8 days after such termination.

9 (B) PROTECTION OF CONFIDENTIAL IN-
10 FORMATION OR TRADE SECRETS.—Before pub-
11 lic disclosure of the documentation under sub-
12 paragraph (A), the Secretary shall delete from
13 the documentation the following:

14 (i) Any information that constitutes
15 trade secret or confidential commercial or
16 financial information.

17 (ii) Any personnel, medical, or similar
18 information, including the serial numbers
19 of implanted devices, which would con-
20 stitute a clearly unwarranted invasion of
21 personal privacy.

1 **SEC. 3. CONDITIONAL CLEARANCE OF CERTAIN MEDICAL**
2 **DEVICES.**

3 (a) IN GENERAL.—Chapter V of the Federal Food,
4 Drug, and Cosmetic Act (21 U.S.C. 351 et seq.) is amend-
5 ed by inserting after section 510 the following:

6 **“SEC. 510A. CONDITIONAL CLEARANCE OF CERTAIN MED-**
7 **ICAL DEVICES.**

8 “(a) IN GENERAL.—Notwithstanding any other pro-
9 vision of law, the Secretary may conditionally clear for in-
10 troduction into interstate commerce for commercial dis-
11 tribution a medical device intended for human use if such
12 medical device is cleared pursuant to section 510(k).

13 “(b) POSTCLEARANCE REQUIREMENTS.—As part of
14 the conditional clearance under subsection (a), the Sec-
15 retary may impose the following:

16 “(1) The Secretary may restrict the sale, dis-
17 tribution, or use of the device but only to the extent
18 that the sale, distribution, or use of the device may
19 be restricted pursuant to section 520(e).

20 “(2) The Secretary—

21 “(A) may require continuing evaluation
22 and periodic reporting on the safety, effective-
23 ness, and reliability of the device for its in-
24 tended use; and

25 “(B) shall, to the extent the Secretary
26 makes a requirement under subparagraph (A),

1 state in the clearance order the reason or pur-
2 pose for such a requirement and the number of
3 patients to be evaluated and the reports re-
4 quired to be submitted.

5 “(3) The Secretary may require a prominent
6 display in the labeling of the device and in the ad-
7 vertising of warnings, hazards, or precautions impor-
8 tant for the device’s safe and effective use, including
9 patient information such as information provided to
10 the patient on alternative modes of therapy and on
11 risks and benefits associated with the use of the de-
12 vice.

13 “(4) The Secretary—

14 “(A) may require maintenance of records
15 that will enable the applicant to submit to the
16 Food and Drug Administration information
17 needed to trace patients if such information is
18 necessary to protect the public health; and

19 “(B) shall, to the extent the Secretary
20 makes the requirement under subparagraph
21 (A), require that the identity of any patient be
22 disclosed in records maintained under the
23 postclearance reporting requirements only to
24 the extent required for the medical welfare of
25 the individual, to determine the safety or effec-

1 tiveness of the device, or to verify a record, re-
2 port, or information submitted to the agency.

3 “(5) The Secretary may require maintenance of
4 records for specified periods of time and organiza-
5 tion and indexing of records into identifiable files to
6 enable the Food and Drug Administration to deter-
7 mine whether there is reasonable assurance of the
8 continued safety and effectiveness of the device.

9 “(6) The Secretary may require submission of
10 periodic reports, at specified intervals, which reports
11 shall comply with the following:

12 “(A) Identify any of the following changes:

13 “(i) New indications for use of the de-
14 vice.

15 “(ii) Labeling changes.

16 “(iii) The use of a different facility or
17 establishment to manufacture, process, or
18 package the device.

19 “(iv) Changes in sterilization proce-
20 dures.

21 “(v) Changes in packaging.

22 “(vi) Changes in the performance or
23 design specifications, circuits, components,
24 ingredients, principle of operation, or phys-
25 ical layout of the device.

1 “(vii) Extension of the expiration date
2 of the device based on data obtained under
3 a new or revised stability or sterility test-
4 ing protocol.

5 “(viii) A change that does not affect
6 the device’s safety or effectiveness.

7 “(B) Contain a summary and bibliography
8 of the following information not previously sub-
9 mitted:

10 “(i) Unpublished reports of data from
11 any clinical investigations or nonclinical
12 laboratory studies involving the device or
13 related devices and known to or that rea-
14 sonably should be known to the applicant.

15 “(ii) Reports in the scientific lit-
16 erature concerning the device and known
17 to or that reasonably should be known to
18 the applicant. If, after reviewing the sum-
19 mary and bibliography, the Food and Drug
20 Administration concludes that the agency
21 needs a copy of the unpublished or pub-
22 lished reports, the Food and Drug Admin-
23 istration shall notify the applicant that
24 copies of such reports should be submitted.

1 “(C) Identify changes made pursuant to an
2 exception or alternative granted under section
3 801.128 or 809.11 of title 21, Code of Federal
4 Regulations.

5 “(7) The Secretary may require batch testing of
6 the device.

7 “(8) The Secretary may provide for any other
8 requirements determined by the Secretary to be nec-
9 essary to provide reasonable assurance, or continued
10 reasonable assurance, of the safety and effectiveness
11 of the device.

12 “(9) The Secretary may require device tracking
13 as provided under part 821 of title 21, Code of Fed-
14 eral Regulations.

15 “(c) RESCISSION OF CONDITIONAL CLEARANCE.—
16 The Secretary may rescind the conditional clearance of a
17 medical device under subsection (a) if the Secretary deter-
18 mines that the conditions imposed on the clearance of the
19 device described in subsection (b) have not been met.”.

20 (b) CIVIL MONETARY PENALTIES.—Section
21 303(f)(1)(A) of the Federal Food, Drug, and Cosmetic Act
22 (21 U.S.C. 333(f)(1)(A)) is amended by inserting “, or
23 a regulation promulgated or an order issued to carry out
24 this Act,” after “any person who violates a requirement
25 of this Act”.

1 (c) PROCESS FOR THE REVIEW OF DEVICE APPLICA-
2 TIONS.—Section 737(8)(J) of the Federal Food, Drug,
3 and Cosmetic Act (21 U.S.C. 379i(8)(J)) is amended by
4 inserting “or required as a condition of clearance of a de-
5 vice under section 510A” after “Act”.

